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RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/192,579A

Input Set : A:\Sequence Listing.txt

Output Set: N:\CRF3\02092001\I192579A.raw

DATE: 02/09/2001 TIME: 14:02:56 RECEIVED

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```
3 <110> APPLICANT: MENOZZI, Franco
            LOCHT, Camille
     4
      6 <120> TITLE OF INVENTION: IDENTIFICATION AND CLONING OF A MYCOBACTERIAL ANTIGEN
             CORRESPONDING TO A HEPARIN-BINDING HAEMAGGLUTININ
     9 <130> FILE REFERENCE: 960-34
    11 <140> CURRENT APPLICATION NUMBER: 09/192,579A
    12 <141> CURRENT FILING DATE: 1998-11-17
    14 <150> PRIOR APPLICATION NUMBER: FR 96 06168
    15 <151> PRIOR FILING DATE: 1996-05-17
    17 <160> NUMBER OF SEQ ID NOS: 20
    19 <170> SOFTWARE: PatentIn Ver. 2.1
    23 <210> SEQ ID NO: 1
    24 <211> LENGTH: 39
    25 <212> TYPE: PRT
    26 <213> ORGANISM: Mycobacterium
    28 <220> FEATURE:
    29 <223> OTHER INFORMATION: sequence comprising a region involved in
             interactions with sulphated glycoconjugates and in
    30
    31
             heparin binding
    33 <400> SEQUENCE: 1
     34 Lys Lys Ala Ala Pro Ala Lys Lys Ala Ala Pro Ala Lys Lys Ala Ala
                                         10
    37 Pro Ala Lys Lys Ala Ala Ala Lys Lys Ala Pro Ala Lys Lys Ala Ala
    38
                   20
                                         25
                                                             30
    40 Ala Lys Lys Val Thr Gln Lys
                35
    41.
    44 <210> SEQ ID NO: 2
    45 <211> LENGTH: 10
    46 <212> TYPE: PRT
    47 <213> ORGANISM: Mycobacterium
    49 <220> FEATURE:
    50 <223> OTHER INFORMATION: peptide S1441
    52 <400> SEQUENCE: 2
    53 Lys Ala Glu Gly Tyr Leu Glu Ala Ala Thr
    58 <210> SEQ ID NO: 3
    59 <211> LENGTH: 18
    60 <212> TYPE: PRT
    61 <213> ORGANISM: Mycobacterium
    63 <220> FEATURE:
    64 <221> NAME/KEY: CDS
    65 <222> LOCATION: (1)
    66 <223> OTHER INFORMATION: peptide S1443; Xaa can be any amino acid
    68 <400> SEQUENCE: 3
W--> 69 Xaa Glu Gly Tyr Val Asp Gln Ala Val Glu Leu Thr Gln Glu Ala Leu
    70 1
    72 Gly Lys
```

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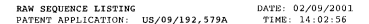
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76 <210> SEQ ID NO: 4
    77 <211> LENGTH: 9
    78 <212> TYPE: PRT
    79 <213> ORGANISM: Mycobacterium
    81 <220> FEATURE:
    82 <221> NAME/KEY: CDS
    83 <222> LOCATION: (1), (4) and (8)
    84 <223> OTHER INFORMATION: peptide S1446; Xaa can be any amino acid
    86 <400> SEQUENCE: 4
W--> 87 Xaa Gln Glu Xaa Leu Pro Glu Xaa Leu
    88 1
    91 <210> SEQ ID NO: 5
    92 <211> LENGTH: 7
    93 <212> TYPE: PRT
    94 <213> ORGANISM: Mycobacterium
    96 <220> FEATURE:
    97 <223> OTHER INFORMATION: Peptide S1447
    99 <400> SEQUENCE: 5
    100 Phe Thr Ala Glu Glu Leu Arg
    101 1
                          5
    104 <210> SEQ ID NO: 6
    105 <211> LENGTH: 17
    106 <21.2> TYPE: DNA
    107 <213> ORGANISM: Mycobacterium
    109 <220> FEATURE:
    110 <223> OTHER INFORMATION: Oligonucleotide originated from the S1441 peptide
    111
              (oligo S1441)
    113 <400> SEQUENCE: 6
                                                                           17
    114 aaggesgagg gstacet
    117 <210> SEQ ID NO: 7
    118 <211> LENGTH: 17
    119 <212> TYPE: DNA
    120 <213> ORGANISM: Mycobacterium
    122 <220> FEATURE:
    123 <223> OTHER INFORMATION: Oligonucleotide originated from the S1441 peptide
              (reverse oligo S1441)
    126 <400> SEQUENCE: 7
    127 aggtascect esgeett
                                                                           17
    130 <21.0> SEQ ID NO: 8
    131 <211> LENGTH: 17
    132 <212> TYPE: DNA
    133 <213> ORGANISM: Mycobacterium
    135 <220> FEATURE:
    136 <223> OTHER INFORMATION: Oligonucleotide originated from the S1443 peptide
    137
              (oligo $1443)
    139 <400> SEQUENCE: 8
                                                                           17
    140 gaccaggesg tsgaget
    143 <210> SEQ ID NO: 9
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144 <211> LENGTH: 17



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145 <212> TYPE: DNA
146 <213> ORGANISM: Mycobacterium
148 <220> FEATURE:
149 <223> OTHER INFORMATION: Oligonucleotide originated from the S1443 peptide
150
          (reverse oligo 51443)
152 <400> SEQUENCE: 9
153 agetesaesg cetggte
                                                                       17
156 <210> SEQ ID NO: 10
157 <211> LENGTH: 21
158 <212> TYPE: DNA
159 <213> ORGANISM: Mycobacterium
161 <220> FEATURE:
162 <223> OTHER INFORMATION: Oligonucleotide named HBHASeq1 and used for
1.63
          sequencing the gene coding for HBHA
165 <400> SEQUENCE: 10
166 agccggtaca acgagctggt c
                                                                       21
169 <210> SEQ ID NO: 11
170 <211> LENGTH: 21
171 <212> TYPE: DNA
172 <213> ORGANISM: Mycobacterium
174 <220> FEATURE:
175 <223> OTHER INFORMATION: Oligonucleotide named HBHA Seqlinv and used for
176
          sequencing the gene coding for HBHA
178 <400> SEQUENCE: 11
179 gaccageteg ttgtacegge t
                                                                       .21
182 <210> SEQ ID NO: 12
183 <211> LENGTH: 19
184 <212> TYPE: DNA
185 <213> ORGANISM: Mycobacterium
187 <220> FEATURE:
188 <223> OTHER INFORMATION: Oligonucleotide named HBHASeq2 and used for
189
          sequencing the gene coding for HBHA
191 <400> SEQUENCE: 12
                                                                       19
192 catecaacae gtegaetee
195 <210> SEQ ID NO: 13
196 <211> LENGTH: 19
197 <212> TYPE: DNA
198 <213> ORGANISM: Mycobacterium
200 <220> FEATURE:
201 <223> OTHER INFORMATION: Oligonucleotide named HBHA Seq3 and used for
202
          sequencing the gene coding for HBHA
204 <400> SEQUENCE: 13
                                                                       19
205 ttgatgtcat caatgttcg
208 <210> SEQ ID NO: 14
209 <211> LENGTH: 19
210 <212> TYPE: DNA
211 <213> ORGANISM: Mycobacterium
213 <220> FEATURE:
214 <223> OTHER INFORMATION: Oligonucleotide named HBHA Seq4 and used for
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```
sequencing the gene coding for HBNA
217 <400> SEQUENCE: 14
218 cgtggaccag gcggtggag
                                                                       19
221 <210> SEQ ID NO: 15
222 <211> LENGTH: 21
223 <212> TYPE: DNA
224 <213> ORGANISM: Mycobacterium
226 <220> FEATURE:
227 <223> OTHER INFORMATION: Oligonucleotide named HBHA Seq 5 and used for
          sequencing the gene coding for HBHA
230 <400> SEQUENCE: 15
231 gacgatcagg aggtttcccc g
                                                                       21
234 <210> SEQ 1D NO: 16
235 <211> LENGTH: 24
236 <212> TYPE: DNA
237 <213> ORGANISM: Mycobacterium
239 <220> FEATURE:
240 <223> OTHER INFORMATION: Oligonucleotide named reverse primer and used for
         sequencing the gene coding for HBHA
243 <400> SEQUENCE: 16
244 agcggataac aatttcacac agga
                                                                       24
247 <210> SEQ ID NO: 17
248 <211> LENGTH: 149
249 <212> TYPE: DNA
250 <213> ORGANISM: Mycobacterium
252 <220> FEATURE:
253 <223> OTHER INFORMATION: nucleotide sequence and amino sequence of a fragment of HBHA
         deduced from a PCR fragment of chromosomal BCG DNA .
254
256 <220> FEATURE:
257 <221> NAME/KEY: CDS
258 <222> LOCATION: (1)..(147)
260 (<400> SEQUENCE: 17
261 aag goo gag ggo tac etc gag goo gog act ago egg tac aac gag etg
                                                                       48
262 Lys Ala Glu Gly Tyr Leu Glu Ala Ala Thr Ser Arg Tyr Asn Glu Leu
                     5
                                         10
                                                             15
265 gtc gag cgc ggt gag gcc gct cta gag cgg ctg cgc agc cag cag agc
266 Val Glu Arg Gly Glu Ala Ala Leu Glu Arg Leu Arg Ser Gln Gln Ser
                20
                                    2.5
267
                                                         3.0
269 tto gag gaa gtg tog gog occ goo gaa ggo tac gtg gac cag gog gto
270 Phe Glu Glu Val Ser Ala Pro Ala Glu Gly Tyr Val Asp Gln Ala Val
            35 .
271
                                 40
273 gag'ct
                                                                       149
274 Głu
277 <210> SEQ ID NO: 18
278 <211> LENGTH: 49
279 <212> TYPE: PRT
280 <213> ORGANISM: Mycobacterium
281 <223> OTHER INFORMATION: amino sequence of a fragment of HBHA deduced from a
286 <400> SEQUENCE: 18
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RAW SEQUENCE LISTING
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TIME: 14:02:56

287 Lys Ala Glu Gly Tyr Leu Glu Ala Ala Thr Ser Arg Tyr Asn Glu Leu 10 290 Val Glu Arg Gly Glu Ala Ala Leu Glu Arg Leu Arg Ser Gln Gln Ser 20 25 30 293 Phe Glu Glu Val Ser Ala Pro Ala Glu Gly Tyr Val Asp Gln Ala Val 40 294 35 45 296 Glu 300 <210> SEQ ID NO: 19 301 <211> LENGTH: 1097 302 <212> TYPE: DNA 303 <213> ORGANISM: Artificial Sequence 305 <220> FEATURE: 306 <223> OTHER INFORMATION: Description of Artificial Sequence: DNA sequence 307 of the BCG gene coding for HBHA 309 <220> FEATURE: 310 <221> NAME/KEY: CDS 311 <222> LOCATION: (331)..(924) 312 <223> OTHER INFORMATION: CDS from 811 to 828, from 829 to 846, from 847 to $864\,,$ from 865 to 885 and from 895 to 915 : peptide 313 314 which may be particularly involved in interactions with sulphated glycoconjugates 317 <400> SEQUENCE: 19 ${\tt 318}\ {\tt cggctggcgg}\ {\tt gtaatcaaac}\ {\tt ctgaaggaca}\ {\tt gtcatctgyg}\ {\tt tgaggtcgac}\ {\tt cgcaggctga}\ {\tt 60}$ 320 tecageegat eggeeggege tggeeaacag egaeteegte gatgaegtge agcaaaggag 120 322 acatgtagtg accggatcag ctgggcctga catchacgaa ctcgaccgac aaccgacccg 180 324 acyatcayga ggtttccccg gcaagtcgcg tgccatgtca atccgcgggt cttgactagt 240 326 cetecetqqa qqaqeegaeg ettgeeccaa egteeaqaee aaagatgtaa gaacgeegat 300 328 atcagaaaat agttaatgaa aggaataccc atg get gaa aac teg aac att gat Met Ala Glu Asn Ser Asn Ile Asp 330 1 332 gac atc aag get eeg ttg ett gee geg ett gga geg gee gae etg gee 333 Asp Ile Lys Ala Pro Leu Leu Ala Ala Leu Gly Ala Ala Asp Leu Ala 334 10 15 20 336 ttg gcc act gtc aac gag ttg atc acg aac ctg cgt gag cgt gcg gag 337 Leu Ala Thr Val Asn Glu Leu Ile Thr Asn Leu Arg Glu Arg Ala Glu 338 25 30 35 340 gag act cgt acg gac acc cgc agc cgg gtc gag gag agc cgt gct cgc 341 Glu Thr Arg Thr Asp Thr Arg Ser Arg Val Glu Glu Ser Arg Ala Arg 342 50 45 344 ctg acc aag ctg cag gaa gat ctg ccc gag cag ctc acc gag ctg cgt 345 Leu Thr Lys Leu Gln Glu Asp Leu Pro Glu Gln Leu Thr Glu Leu Arg 65 348 gag aag ttc acc gcc gag gag ctg cgt aag gcc gcc gag ggc tac ctc 594 349 Glu Lys Phe Thr Ala Glu Glu Leu Arg Lys Ala Ala Glu Gly Tyr Leu 75 80 85 352 gag gee geg act age egg tac aac gag etg gte gag ege ggt gag gee 353 Glu Ala Ala Thr Ser Arg Tyr Asn Glu Leu Val Glu Arg Gly Glu Ala 354 90 95 100 356 gct cta gag cgg ctg cgc agc cag cag agc ttc gag gaa gtg tcg gcg

 VERIFICATION SUMMARY
 DATE: 02/09/2001

 PATENT APPLICATION:
 US/09/192,579A
 TIME: 14:02:57

Input Set : A:\Sequence Listing.txt
Output Set: N:\CRF3\02092001\I192579A.raw

L:69 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:3 L:87 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:4